

UNITED STATES ENVIRONMENTAL PROTECTION AGENCY WASHINGTON, D.C. 20460

OCT A 1989

OFFICE OF
SOLID WASTE AND EMERGENCY RESPONSE

TO ALL NRC LICENSEES:

SUBJECT: GUIDANCE ON THE DEFINITION AND IDENTIFICATION OF

COMMERCIAL MIXED LOW-LEVEL RADIOACTIVE AND HAZARDOUS

WASTE AND ANSWERS TO ANTICIPATED QUESTIONS

The U.S. Environmental Protection Agency (EPA) has jurisdiction under the Resource Conservation and Recovery Act (RCRA) over the management of wastes with the exception of radioactive wastes subject to the Atomic Energy Act (AEA). Accordingly, commercial use and disposal of source, byproduct and special nuclear material wastes are regulated by the U.S. Nuclear Regulatory Commission (NRC) to meet EPA environmental standards. Under the AEA Low-Level Radioactive Wastes (LLW) contain source, byproduct, or special nuclear material, but they may also contain chemical constituents which are hazardous under EPA regulations in 40 CFR Part 261. Such wastes are commonly referred to as Mixed Low-Level Radioactive and Hazardous Waste (Mixed LLW).

NRC regulations exist to control the byproduct, source, and special nuclear material components of commercial Mixed LLW; EPA has the authority and continues to develop regulations to control the non-radioactive component of the Mixed LLW. Thus, the individual constituents of commercial Mixed LLW are subject to either NRC or EPA regulations. However, when the components are combined to become Mixed LLW, neither statute has exclusive jurisdiction. This has resulted in a situation of dual regulation where both NRC and EPA may regulate the same waste.

Enclosed is the revised guidance document entitled, "Guidance on the Definition and Identification of Commercial Mixed Low-Level Radioactive and Hazardous Waste." This document was developed jointly by the NRC and EPA to aid commercial LLW generators in assessing whether they are currently generating Mixed LLW. It is based on NRC and EPA regulations in effect on December 31, 1988.

Notice of availability of for comments were published in 1987, and comments were subsequible comment in the question document to provide clarificative raised.

GUIDANCE ON THE DEFINITION AND IDENTIFICATION OF COMMERCIAL MIXED LOW-LEVEL RADIOACTIVE AND HAZARDOUS WASTE

Definition

Mixed Low-Level Radioactive and Hazardous Waste (Mixed LLW) is defined as waste that satisfies the definition of low-level radioactive waste (LLW) in the Low-Level Radioactive Waste Policy Amendments Act of 1985 (LLRWPAA) and contains hazardous waste that either (1) is listed as a hazardous waste in Subpart D of 40 CFR Part 261 or (2) cause the LLW to exhibit any of the hazardous waste characteristics identified in Subpart C of 40 CFR Part 261.

Identification

The policy provided in this guidance was developed jointly by the U.S. Nuclear Regulatory Commission (NRC) and the U.S. Environmental Protection Agency (EPA). LLW that contains hazardous wastes defined under the Resource Conservation and Recovery Act (RCRA) is Mixed LLW. Under current Federal law, such waste is subject to regulation by NRC under the Atomic Energy Act (AEA), as amended, and by EPA under RCRA, as amended. In the absence of legislation to the contrary, management and disposal of this waste must be conducted in compliance with NRC and EPA or equivalent state regulations.

This guidance presents a methodology (Figure 1) that may be used by generators of commercial LLW to identify Mixed LLW. Implementation of the methodology should identify Mixed LLW and aid generators in assessing whether they are currently generating Mixed LLW. Generators are cautioned, however, that application of the methodology does not affect the need to comply with applicable NRC and EPA regulations. Because EPA's regulations for hazardous waste are currently changing, generators should use applicable regulations that are in effect at the time of implementation of the methodology. This guidance has been prepared based on NRC and EPA regulations in effect on December 31, 1988.

Application of this methodology to identify Mixed LLW will reveal the complexities of the definition of Mixed LLW. If generators have specific questions about whether LLW is Mixed LLW, they should promptly contact the agencies by writing to the persons listed below. For questions about whether the waste is low-level radioactive waste, contact:

Mr. Dan E. Martin
Division of Low-Level Waste
Management and Decommissioning
U.S. Nuclear Regulatory Commission
Mail Stop WF5E4
Washington, D.C. 20555

For questions about whether the waste is hazardous waste, contact:

Ms. Betty Shackleford
Mixed Waste Coordinator
Permits and State
Programs Division
Mail Code OS-342
U.S. Environmental
Protection Agency
401 M St., S.W.
Washington, D.C. 20460

Methodology

Step 1. Identify LLW

Step 1 in the methodology requires that the generator determine whether the waste is LLW as defined in the LLRWPAA. This Act defines LLW as radioactive material that (A) is not high-level radioactive waste, spent nuclear fuel, or byproduct material as defined in section 11e(2) of the AEA (i.e., uranium or thorium mill tailings) and (B) the NRC classifies as LLW consistent with existing law and in accordance with (A). If the generator determines that the waste is LLW, the generator should proceed to step 2. If the determination is negative, then the waste cannot be Mixed LLW because it is not LLW. However, the waste may be another radioactive or hazardous waste regulated under AEA, RCRA, or both statutes.

Step 2. Identify Listed Hazardous Waste

In step 2, the generator determines whether the LLW contains any hazardous wastes listed in Subpart D of 40 CFR Part 261. Subpart D of Part 261 is reproduced in Appendix I of this guidance. LLW is Mixed LLW if it contains any hazardous wastes specifically listed in Subpart D of 40 CFR Part 261. Listed hazardous wastes include hazardous waste streams from specific and non-specific sources listed in 40 CFR Parts 261.31 and 261.32 and discarded commercial chemical products listed in 40 CFR Part 261.33. The generator is responsible for determining whether LLW contains listed hazardous wastes. The determination should be based on knowledge of the process that generates the waste. For example, if a process produces LLW that contains spent solvents that are specifically listed in the tables of Subpart D of Part 261, the generator should suspect that the waste is Mixed LLW.

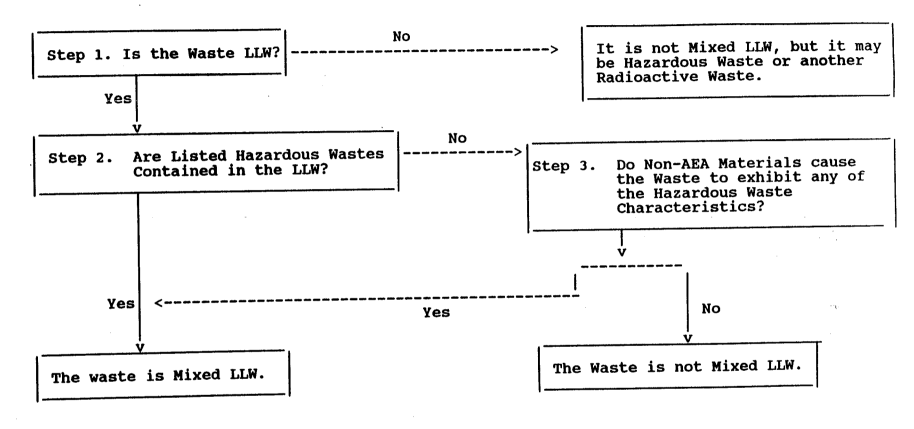


Figure 1. Identification of Mixed LLW.

Step 3. Identify Hazardous Characteristics

If the LLW does not contain a listed hazardous waste, Step 3 of the methodology requires the generator to determine whether the LLW contains hazardous wastes that cause the LLW to exhibit any of the hazardous waste characteristics identified in Subpart C of 40 CFR Part 261. This determination can be based on either (1) an assessment of whether the LLW exhibits one or more of the hazardous waste characteristics because it contains non-AEA materials (i.e., materials other than source, special nuclear, and byproduct materials) based on the generator's knowledge of the materials or processes used in generating the LLW or (2) testing of the LLW in accordance with the methods identified in Subpart C of Part 261. Except for certain ores containing source material, which are defined as source material in 10 CFR 40.4(h), and uranium and thorium mill tailings or wastes, NRC and EPA interpret the definitions of source, special nuclear, and byproduct materials to include only the radioactive elements themselves. Generators should identify non-AEA materials contained in the LLW by examining the process that generates the waste. For example, if the process mixes byproduct material (an AEA material) with a volatile organic solvent (a non-AEA material), the generator would determine either through his knowledge or testing of representative samples of the LLW that contain the solvent waste whether the waste exhibits any of the hazardous waste characteristics because it contains the solvent.

If the wastes are tested, the generator should collect and test representative samples of the LLW to determine if the waste exhibits any of the characteristics identified in Subpart C because it contains the non-AEA materials. These characteristics include ignitability (Section 261.21), corrosivity (Section 261.22), reactivity (Section 261.23), and Extraction Procedure (EP) toxicity (Section 261.24). Waste testing should be conducted in a manner that is consistent with the worker protection requirements in 10 CFR Part 20. The purpose of the characteristics tests is to identify hazardous wastes that are not specifically listed in Subpart D of 40 CFR Part 261. Test methods to collect representative samples of wastes are described in Appendix I of 40 CFR Part 261. The samples should then be tested using the referenced testing protocols (e.g., ASTM Standard D-93-79 or D-93-80 for the Pensky-Martens Closed Cup Ignitability Test). EPA's testing requirements are reproduced in Appendix II of this guidance. It should be noted that on June 13, 1986, EPA proposed a modification to the EP Toxicity testing requirements to include organic constituents.

If LLW contains a listed hazardous waste or non-AEA materials that cause the LLW to exhibit any of the hazardous waste characteristics, the waste is Mixed LLW and must, therefore, be managed and disposed of in compliance with EPA's Subtitle C hazardous waste regulations in 40 CFR Parts 124, and 260 through 270, and NRC's regulations in 10 CFR Parts 20, 30, 40, 61, and 70.

Management and disposal of Mixed LLW must be conducted in compliance with state requirements in states with EPA-authorized regulatory programs for the hazardous components of such waste and NRC agreement state radiation control programs for LLW.

Questions and Answers

As a supplement to the Guidance on the Definition and Identification of Commercial Mixed Low-Level Radioactive and Hazardous Waste (Mixed LLW), answers to anticipated questions are included to clarify obscure points and to respond to public comments.

1. Are my low-level radioactive wastes exempt from RCRA because they are source, special nuclear, or byproduct materials as defined under the AEA?

Except for certain ores containing source material, which are defined as source material in 10 CFR 40.4(h), and uranium and thorium mill tailings or wastes, NRC and EPA consider that only the radionuclides themselves are exempt from RCRA. Section 1004(27) of RCRA excludes source, special nuclear, and byproduct material from the definition of "solid waste." RCRA defines solid waste as:

"any garbage, refuse, sludge from a waste treatment plant, water supply treatment plant, or air pollution control facility and other discarded material, including solid, liquid, semisolid, or contained gaseous material resulting from industrial, commercial, mining, and agricultural operations, or from community activities, but does not include solid or dissolved materials in irrigation return flows or industrial discharges which are point sources subject to permits under section 402 of the Federal Waster Pollution Control Act, as amended (86 Stat. 880), or source, special nuclear, or byproduct material as defined by the Atomic Energy Act of 1954, as amended (68 Stat. 923)." [emphasis added]

Since "hazardous waste" is a subset of "solid waste," RCRA also excludes source, special nuclear, and byproduct materials from the definition of hazardous waste and, therefore, from regulation under EPA's RCRA Subtitle C program. Section 11 of the Atomic Energy Act, as amended, defines these radioactive materials as follows:

Source material means (1) uranium, thorium, or any other material which is determined by the Atomic Energy Commission (AEC) pursuant to the provisions of section 61 of the AEA to be source material, or (2) ores containing one or more of the foregoing materials, in such concentration as the AEC may by regulation determine from time to time.

Special nuclear material means (1) plutonium, uranium enriched in the isotope 233 or in the isotope 235, and any other material which the AEC, pursuant to the provisions of Section 51 of the AEA, determines to be special nuclear material; or (2) any material artificially enriched by any of the foregoing, but does not include source material.

Byproduct material means (1) any radioactive material (except special nuclear material) yielded in or made radioactive by exposure to radiation incident to the process of producing or utilizing special nuclear material, and (2) the tailings or wastes produced by the extraction or concentration of uranium or thorium from any ore processed primarily for its source material content.

Source, special nuclear, and byproduct materials, however, may be mixed with other radioactive or non-radioactive materials that are not source, special nuclear, or byproduct materials. For example, tritium may be contained in toluene, a nonhalogenated aromatic solvent. Consistent with the definition of byproduct material, the tritium may be considered a byproduct material, while the toluene that contains the tritium would not be byproduct material. Mixtures of toluene and tritium could satisfy the definition of Mixed LLW because they contain listed hazardous waste (spent toluene) and tritium that may qualify as LLW if it has been produced by activities regulated by NRC under the AEA.

2. What are some examples of Mixed LLW?

A preliminary survey performed for the NRC identified two potential types of Mixed LLW:

- o LLW containing organic liquids, such as scintillation liquids and vials; organic lab liquids; sludges; and cleaning, degreasing, and miscellaneous solvents.
- o LLW containing heavy metals, such as discarded lead shielding, discarded lined containers, and lead oxide dross containing uranium oxide; light water reactor (LWR) process wastes containing chromate and LWR decontamination resins containing chromium; and mercury amalgam in trash.

The preliminary survey concluded that potential Mixed LLW comprises a small percentage of all LLW. For example, LLW containing organic liquids accounted for approximately 2.3% by volume of LLW reported in the preliminary survey (Bowerman, et al., 1985).

An earlier survey identified a more diverse universe of potential Mixed LLW including wastes that contained aldehydes, aliphatic halogenated hydrocarbons, alkanes, alkenes, amino acids, aromatic hydrocarbons, chelating agents, esters, ethers, ketones, nitrosamines, nucleotides, pesticides, phenolic compounds, purines, resins, steroids, and vitamins (General Research Corporation, 1980). NRC also anticipates that additional LLW may be identified as Mixed LLW in the future, as generators implement the definition of Mixed LLW and as EPA revises the definition of hazardous waste.

3. Could some "below regulatory concern" wastes be considered Mixed LLW?

A determination that radioactive wastes are below regulatory concern (BRC) for radioactivity may affect how the wastes are managed or discarded, but it does not affect the legal status of the wastes. Specifically, their status with respect to the definition of Mixed LLW does not change. BRC waste is still LLW because it satisfies the definition of LLW in the LLRWPAA and is within the NRC's jurisdictional authority under the AEA.

When radioactive waste contains sufficiently low concentrations or quantities of radionuclides, NRC may find that they do not need to be managed or disposed of as radioactive wastes. For NRC to make such a finding, management and disposal of the waste must not pose an undue radiological risk to the public and the environment. However, NRC's determination that the radioactive content of the wastes is below NRC regulatory concern does not relieve licensees from compliance with applicable rules of other agencies governing non-radiological hazards (e.g., regulations of EPA or the Department of Transportation).

Therefore, some BRC wastes may still be considered Mixed LLW if they contain hazardous wastes that have been listed in Subpart D of 40 CFR Part 261 or that cause the LLW to exhibit any of the hazardous characteristics described in Subpart C of 40 CFR Part 261. BRC Mixed LLW may be managed without regard to its radioactivity (but it must still be managed as a hazardous waste in compliance with EPA's regulations for hazardous waste generation, storage, transportation, treatment, and disposal (cf. 40 CFR Parts 262 through 266)).

4. If I use chemicals in my process that are identified by EPA as hazardous constituents, should I assume that my LLW is Mixed LLW?

No. Low-level radioactive waste that contains hazardous constituents may not necessarily be Mixed LLW. As defined above, Mixed LLW is LLW that contains a known hazardous waste (i.e., a listed hazardous waste) or that exhibits one or more of the hazardous characteristics because it contains non-AEA materials. For wastes that are not listed in Subpart D of 40 CFR Part 261, testing is not necessarily required to "determine" whether the LLW exhibits any of the hazardous characteristics. A generator may be able to determine whether the LLW is Mixed LLW based on knowledge of the waste characteristics or the process that generates the LLW.

Furthermore, if the generator normally segregates LLW from hazardous and other types of wastes, there is no need to assume that hazardous wastes may have been inadvertently mixed with LLW or to inspect each container or receptacle to ensure that inadvertent mixing has not occurred. Although the generator is subject to RCRA inspections and must follow the manifest, pre-transport, and other requirements of

40 CFR Part 262, the generator is not required to demonstrate that every LLW container does not contain hazardous waste.

5. How can I obtain representative samples of heterogeneous trash included in LLW to perform the hazardous characteristics tests?

Before discussing the collection of representative samples of waste, generators are reminded that they are not required to test LLW to determine if the waste contains hazardous wastes. Generators and handlers of mixed waste and hazardous waste can declare their wastes hazardous or nonhazardous based on knowledge of the process/production of the waste, in lieu of testing for a characteristic.

Representative samples of waste should be collected for testing in accordance with EPA's regulations in 40 CFR 261.20(c), which state that waste samples collected using applicable methods specified in Appendix I of Part 261 will be considered as representative samples for hazardous characteristics testing. This appendix has been included in its entirety in Appendix II of this guidance. The sampling techniques described in Appendix I of Part 261 apply to extremely viscous liquids, fly ash-like material, containerized liquid wastes, and liquid wastes in pits, ponds, lagoons, and similar reservoirs. In the absence of guidance about sampling heterogeneous wastes, generators should use appropriate portions of the sampling methods described in Appendix I of Part 261 and EPA's manual entitled "Test Methods for Evaluating Solid Waste, Third Edition (i.e., SW-846) in combination with other methods to collect, to the maximum extent practicable, representative samples of the waste to be tested.

6. Are lead containers whose primary use is for shielding in disposal operations, hazardous waste under RCRA?

No. While lead containers and lead container liners may exhibit the hazardous characteristic for lead, those containers whose primary use is for shielding in low-level waste disposal operations are not considered wastes and thus, are not subject to the hazardous waste rules. These same containers and liners if disposed of or discarded would be considered wastes and if they exhibit the hazardous characteristic, would be subject to the hazardous waste rules.

It should be noted that EPA recognizes that all lead containers and liners may be equally hazardous to human health and the environment when placed in the ground independent of its legal classification as a waste or container. Therefore, EPA recommends that all lead containers and lead liners be managed in an environmentally safe manner (e.g., managed in a permitted hazardous waste facility or treated such that it no longer exhibits its characteristic).

Encapsulation may be a viable mechanism to mitigate lead migration from these containers and liners. The EPA has not evaluated specific containers or encapsulation methodologies using the EP Toxicity test.

7. If a waste contains any of the constituents listed on Appendix VIII of Part 261, is it a hazardous under RCRA?

No. Under RCRA, a waste is hazardous if it is a "listed" waste or it exhibits a hazardous characteristic. Wastes are listed by EPA if they contain significant amounts of toxic constituents identified in Appendix VIII, and the Agency has determined that these toxic constituents are persistent and mobile to some degree such that they pose a potential and substantial threat to human health and the environment. (Factors outlined in 40 CFR 261.11(a)(3)(i)-(xi), which include nature of the toxicity present and potential degradation products, may be considered when determining whether or not a waste should be listed). However, until the Agency lists the wastes in Subpart D of Part 261, they would not be considered hazardous by EPA (even if the waste contains one or more of the hazardous constituents listed on Appendix VIII) unless the waste would exhibit one or more of the hazardous waste characteristics.

References

- Bowerman, B. S., Kempf, C. R., MacKenzie, D. R., Siskind, B. and P. L. Piciulo, 1985, "An Analysis of Low-Level Wastes: Review of Hazardous Waste Regulations and Identification of Radioactive Mixed Wastes," NUREG/CR-4406, U.S. Nuclear Regulatory Commission.
- General Research Corporation, 1980, "Study of Chemical Toxicity of Low-Level Wastes," NUREG/CR-1793, U.S. Nuclear Regulatory Commission.

Appendix I

Subpart D—Lists of Hezardous Wastes

£ 261.30 General.

(a) A solid waste is a hazardous waste if it is listed in this subpart, unless it has been excluded from this list under \$\frac{1}{2}\$ 260.20 and 260.22.

(b) The Administrator will indicate

his basis for listing the classes or types of wastes listed in this Subpart by employing one or more of the following Hazard Codes:

Ignisble Waste	(1
F-3-4-5046 At State	Ć
Corroene Waste	滴
Recove Waste	Œ
EP Tosc Waste	Ä
Acute Hexardous Waste	'n
Total Waster	• • •

Appendix VII identifies the constituent which caused the Administrator to list the waste as an EP Toxic Waste (E) or Toxic Waste (T) in §§ 261.31 and 261.32.

(c) Each hazardous waste listed in this subpart is assigned an EPA Hazardous Waste Number which precedes the name of the waste. This number must be used in complying with the notification requirements of Section 3010 of the Act and certain record-keeping and reporting requirements under Parts 262 through 255 and Part 270 of this chapter.

(d) The following hazardous wastes listed in § 261.31 or § 261.32 are subject to the exclusion limits for acutely hazardous wastes established in § 261.5: EPA Hazardous Wastes Nos. FO20. FO21, FO22, FO23, FO26, and FO27.

[45 FR 33119, May 19, 1980, as amended at 48 FR 14294, Apr. 1, 1983; 80 FR 2000, Jan. 14, 1985]

The following solid wastes are listed hazardous wastes from non-specific sources unless they are excluded under \$\$ 260.20 and 260.22 and listed in Appendix IX.

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Menc	:		
÷oc•	The independ energy horsespecial expenses and in section 7	_	
	The following spent happenated solvents used in pagressing. Tetrachiorpethylene shorted in the trachiorpethylene carpon retrachionse.		
	and chionneted fluorocarbons, all spent solvent motures/biends used in pagrasa-		
	ing combaning, before use a total of ten percent or more liby volumes of one or		
	more of the above hatogenated solvents or those solvents rated in F002 F004 and F005, and self-bottoms from the recovery of these spent solvents and solent		
	sowers, matures.		
•	The following sperit helogenated solvents: "etrachlorostiyiene: metriviene chionos		
	trichlorosthytene. 1.1.1-trichlorosthene chlorobertzene. 1.1.2-trichloro-		
	cetteris, critic-dichicrobenzene stichicrofluoromethane, and 1.1.2-trichicroethane all spent solvent monunes/blends containing, before use, a lotar of ten percent or		
	more (by volume) of one or more of the above halogenesed solvents or those		
	letted in F001 F004, or F005, and still bottome from the recovery of these spent		
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	metherios at spent solvent matures/blends containing, before use ignly the apove- spent non-halogenisted solvents, and all spent solvent matures/blends containing.		
	before use, one or more of the above non-halogenesed solvents, and, a total of		
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	retropersions, all spent envent matures/blands containing, before use, a total of	1,	
	ten parcent or more (by volume) of one or more of the above non-halogenated		
	solvents or those solvents issed in F001 F002, and F005 and sell bottoms from		
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² 205	The lollowing spent non-harogeneted solvents. Toluene, methyl ethyl xetone, carbon daufilds, exbutanol, pyrigine bergene 2-ethoxyethanol, and 2-intropropere, all	0.23	
	spent solvent mixtures/biends containing, before use, a total of len percent or		
	more (by volume) of one or more of the above non-halogeneted solvents or those		
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•••	these spent solvents and spent solvent textures	_	
*206	Wastewater treatment sudges from electroplating operations except from Tie	in	
	following processes, (1) Surfunc acid endoting of eluminum (2) tin pleting on carbon steel (3) zinc pleting (segregated beels) on carbon steel (4) aluminum or		
	and-elumenum pleang on certain steel (5) cleaning/strating electrosed with an		
	and and aluminum pleang on cardon steel, and (6) chemical eldring and milling of		
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	CYSNIGOS are used in the process		
±010	Quenching both readures from on boths from motel heat treating operations where treatment are used in the process.	(* H	
5011	Spent cyanate solutions from set beth pot clearing from metal feet treating	(R T)	
	CONTROLS.		
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(46 PR 4617, Jan. 16, 1961, as amended at 46 PR 27477, May 20, 1961; 49 PR 5312, Peb. 10, 1964; 49 PR 37070, Sept. 21, 1964; 50 PR 665, Jan. 4, 1965; 50 PR 2000, Jan. 14, 1965, 50 PR 53319, Dec. 31, 1965, 51 PR 2702, Jan. 21, 1966; 51 PR 6541, Peb. 25, 1986)

\$251.32 Hazardous wastes from specific sources.

The following solid wastes are listed hazardous wastes from specific sources unless they are excluded under §§ 260.20 and 260.22 and listed in Appendix IX.

Industry and EPA hezardous waste No.	Hezardous weste	Heanro 6004
Wood preserveson: K001	Bottom sediment studge from the treatment of wastewaters from wood preserving processes that use creosole and/or pentachlorophenol.	e
kooz	Wassewater treatment studge from the production of chrome yellow and orange asoments.	Э
K003	Wastewater treatment studge from the production of molybdate orange pigments	Э
K004	Wastewater treatment sludge from the production of zinc yellow pigments	Œ
K005	Wastewater treatment studge from the production of chrome green pigments	E
K006	Wassewater treatment sludge from the production of officeme code green pigments	e
NOO	(anhydrous and hydrated)	
K007	Wastewater treatment studge from the production of iron blue pigments	m
K008	Oven residue from the production of ahrome ciede green pigments	m
Organic chemicals:		۱
K009	Distillation bottoms from the production of acetalde/tyde from ethylene	m .
K010	Dublishon side cuts from the production of agestidefivide from ethylene	וש_
K011	Bottom streem from the wastewater stropper in the production of acryloneme	(A, T)
K013	Bottom stream from the acetoninie column in the production of scrytoninis	(P, T)
K014	Bottoms from the acetonimie puniceson column in the production of acrytomitie	m .
K015	Still bottoms from the destilation of benzyl chionds	<u>m</u>
K016	Heavy ends or distillation residues from the production of carbon tetrachlonds	(n)
K017	Heavy ends (self bottoms) from the purification enturn in the production of	(m
	epichlorohydnn.	m
K018	Heavy ends from the tractionation column in ethyl chloride production	i iii
K019	Meany erics from the distriction of employs dictionals in employee dictionals graduction.	i '''
K020	Heavy ands from the destination of veryl otherise in veryl chloride monomer	TO .
	production. Aqueous scent aritmony catalyst waste from fluoremethenes production	m
K021	Agusous sport aremony complete water from adoptive transport from California	lκ
K0:22	Distillation bodom tare from the production of phensi/apstone from currents	m
K023	Detriction bottoms from the production of princing annythree from rephthetiens] iii
K024	Dispersion porjous sour sus buggings of business sustained anniverse sourcement	li
KC93	Desileton light ends from the production of phthalic anhydrids from ortho-sylene	lω
K094	Distillation bottoms from the production of prichatic anhydride from antitio-nylene] iii
K025	Distillation bottoms from the production of nitrobangens by the nitration of bungane	liii
K026	Stroping self tests from the production of mothy othyl pyrianes	lia n
K027	Continue and detriation residues from toluene discoyensie production	in "
K029	Spent except from the hydrachionnetor reactor in the production of 1,1,1-monlor- cethere.	1
K029	Waste from the product steem stroper in the production of 1,1,1-transcriptions	ወ
K005	Distriction bottoms from the production of 1,1,1-but mercethere	<u>−</u>
K004	Meany ands from the heavy ands estumn from the production of 1,1,1-incherostitues.	m
Y.000	Column bottoms or heavy ends from the serrained production of trishteresthylene	æ
	and percharacthylene.	_ m
K083	Distillation bottoms from aniline production	1 ന്
K103	Process residues from entire extraction from the preduction of entire	1 iii
K104	Commence westinister strooms generated from introduction arrangement production	1 iii
K085	Distillation or fractionation column bottoms from the preduction of distributions	186
K106	Separated Squares stream from the reactor product wearing step in the production of phorobensenes.	'''
Inorganic ehemicals:		1_
K071	Brine purification mude from the mereury cell process in charmo production, where	(n
	accurately greaterfied inner at not used.	1_
X073	Channeled hydroperton waste from the sufficient step of the deshroom cell	m
·-·	archine using graphic angles in chienne production.	1_
K106	Washpuster treatment studge from the mereury cell process in criterine preduction	⊣m
Postcries:	· •	1_
K031	By-product solia generated in the production of MDMA and escendylic acid	ዛ መ
K032	Manhanatar trackment charles from the small track of Carlot Tolling to the Carlot Tollin	-l Œ
K033	" Managerates and occup water that are quantities on chambers as an	i in
	1 and mos of chicologs	
10034	Piler saids from the Strazen of househiorecyclepentadiene in the production of	(m
	ensertene Vascum straper dechargo from the attendanc attendant in the production of	m
· 14397	- Animana	
10094	Manager transport student experient in the studention of process	_ ლ
10006		-1 ©
10037		<u>اس</u>
10036	Washington from the washing and straining of phorse production. Filter case from the Straten of distriptionspheredities and in the production of	-Jm
1000	Filter sets truth the filtraten of destrutemental states and in the production of	(III)
*****	anaria.	
10040	Wassender segment studge from the production of phoreto	⊣ <u>m</u>
KOM1	Wasserster Presmert studies from the production of temphone	<u> ქლ</u>
1000	Untrasted process wastewater from the production of temphone	ന

industry and EPA hazardous waste No.	Hezardous waste		
KG42	Heavy ends or distillation reactures from the distillation of tetrachlorobertzene in the production of 2.4.5-T.	m	
K043	2 & Dehistography waste from the production of 2.4-0	Θ.	
K099	Untracted westewater from the production of 2.4-D	9	
Evolutives:		1	
K044	Wastewater treatment sludges from the manufacturing and processing of explosives	(P)	
K045	Scent carbon from the treatment of westewater containing explosives	(PI)	
K046	Wassewater treatment studges from the menufacturing, formulation and loading of lead-based shoeting compounds.	Э	
K047	Pint/red water from TNT operations	(PI)	
Petroleum refinanc:		i ' '	
K048	Descrived at floration (DAF) float from the petroleum refining industry	m	
K049	Sign oil emutsion solids from the petroleum refining industry	m	
K050	Heat exchanger bundle cleaning studge from the petroleum refining industry	m	
K051	AP! separator studge from the petroloum refining industry	m	
KOA2	Tank bottoms (loaded) from the perclaum refining industry	m	
iven and steel:		1 ' '	
K061	Emission control dust/studge from the primary production of steel in electric	Œ	
K062	Spent proble Equar generated by stool finishing operations of plants that produce iron or steel.	EC.T;	
Secondary lead:		١	
K009	Emeson control dust/studge from secondary lead smelting	ת ו	
K100	Waste leaching solution from acid leaching of emission eartifol dust/studge from secondary tead smalling.	LG.	
Vetennery phermaceutosis:	1	i_	
K084	Wasseveter treatment studges generated during the production of veterinary pharme- causesis from arsenic or organo-arsenic compounds.	m	
K101	Dissiliston for residues from the distillation of antine-based compounds in the production of veterinary pharmacoulocals from areanic or engano-areanic compounds.	m	
K102	Residue from the use of activated carbon for decolorization in the graduction of veterinary phyrmacoulocals from arsenic or organo-arsenic compounds.	m	
ink formulation: K086	Solvent washes and studges, causic washes and studges, or wash washes and studges from cleaning tups and equipment used in the formulation of this from pigments, dnors, soeps, and stabilizers containing chromium and lead.	Э	
Colung:		1_	
K060	Ammons self time studge from coking operations	<u>.</u> m	
K067	Decemer tens ter studge from cotting operations.	ıπ	

[46 FR 4618, Jan. 16, 1981, as amended at 46 FR 27476-27477, May 20, 1981; 49 FR 27070, Sept. 21, 1984; 50 FR 42942, Oct. 23, 1985; 51 FR 5330, Feb. 13, 1986; 51 FR 19322, May 28, 1986]

EFFECTIVE DATE NOTE At 51 FR 5330, Peb. 13, 1985, in § 261.32, waste streams "K117, K118, and K136" in the subgroup "Organic Chemicals" were added, effective August 13, 1986.

\$ 251.33 Discarded commercial chemical products, off-specification species, container residues, and spill residues thereof.

The following materials or items are hazardous wastes if and when they are discarded or intended to be discarded, when they are mixed with waste oil or used oil or other material and applied to the land for dust suppression or road treatment, or when, in lieu of their original intended use, they are produced for use as (or as a component of) a fuel, distributed for use as a fuel.

(a) Any commercial chemical product, or manufacturing chemical intermediate having the generic name listed in paragraph (e) or (f) of this section.

- (b) Any off-specification commercial chemical product or manufacturing chemical intermediate which, if it met specifications, would have the generic name listed in paragraph (e) or (f) of this section.
- (c) Any container or inner liner removed from a container that has been used to hold any commercial chemical product or manufacturing chemical intermediate having the generic names listed in paragraph (e) of this section, or any container or inner liner removed from a container that has been

used to hold any off-specification chemical product and manufacturing chemical intermediate which, if it met specifications, would have the generic name listed in paragraph (e) of this section, unless the container is empty as defined in § 261.7(b)(3) of this chapter.

IComment: Unless the residue is being beneficially used or reused, or legitimately recycled or reclaimed: or being accumulated, stored, transported or treated prior to such use, re-use, recycling or reclamation. EPA considers the residue to be intended for discard, and thus a hazardous waste. An example of a legitimate re-use of the residue would be where the residue remains in the container and the container is used to hold the same commercial chemical product or manufacturing chemical product or manufacturing chemical intermediate it previously held. An example of the discard of the residue would be where the drum is sent to a drum reconditioner who reconditions the drum but discards the residue.]

(d) Any residue or contaminated soil, water or other debris resulting from the cleanup of a spill into or on any land or water of any commercial chemical product or manufacturing chemical intermediate having the generic name listed in paragraph (e) or (f) of this section, or any residue or contaminated soil, water or other debris resulting from the cleanup of a spill, into or on any land or water, of any off-specification chemical product and manufacturing chemical intermediate which, if it met specifications, would have the generic name listed in paragraph (e) or (f) of this section.

[Comment: The phrase "commercial chemical product or manufacturing chemical intermediate having the generic name listed in . . ." refers to a chemical substance which is manufactured or formulated for commercial or manufacturing use which consists of the commercially pure grade of the chemical, any technical grades of the chemical that are produced or marketed, and all formulations in which the chemical is the sole active ingredient. It does not refer to a material, such as a manufacturing process waste, that contains any of the substances listed in paragraph (e) or (f). Where a manufacturing process waste is deemed to be a hazardous waste because it contains a substance listed in paragraph (e) or (f), such waste will be listed in either § 251.31 or § 251.32 or will be identified as a hazardous waste by the characteristics set forth in Subpart C of this part.)

(e) The commercial chemical products, manufacturing chemical intermediates or off-specification commercial chemical products or manufacturing chemical intermediates referred to in paragraphs (a) through (d) of this section, are identified as acute hazardous wastes (H) and are subject to be the small quantity exclusion defined in § 261.5(e).

(Comment: For the convenience of the regulated community the primary hazardous properties of these materials have been indicated by the letters T (Toxicity), and R (Reactivity). Absence of a letter indicates that the compound only is listed for acute toxicity.]

These wastes and their corresponding EPA Hazardous Waste Numbers are:

Haserdous waste No.	
PORS	Asstaldehyde, shipro-
POGS	Acsternide, N-Commothicsomothy-Q-
P067	Asstarrate, 2-Busto-
P066	Acres and, flore, andus and
P086	Assumide and N-Emetrylan-
	(templicity)this, matryl exter
P001	3-(alpha-Acatery/barzy0-4-hydronysta/marin
	Und sale, when present at exhaustrations
	(Franker Share 0.3%
P002	1-Acroyl-2-Diseases
P003	Agration
P070	Alderb
P004	Aleto
P006	Allyl alcohol
P006	Aluminum phosphide S-(Ameromethyl)-3-isosphid
P000	4.04
P008	4-pAmospyraine Ammerium pistate (ft)
P118	Artempreum venedate
P010	Arcents said
P012	America (NII) saide
P011	Argenic (V) enide
P011	Argente pentende
P012	Argents Wester
P036	Arene, dethyl-
P064	Asridine
P013	Berlum oyanida
P024	Beresnamme, 4-eNoro-
P077	Bersenemine, 4-ratio-
P029	Bersone, (chloremethyl)-
P042	12-Sermenadol, 4-(1-hydrany-2-methyl-
P014	Orninalativi)- Bermanusiusi
PO26	Bench entendo
PO15	Boryoum dust
POIA	Blackteremethyl ether
P017	Brumeastene
P018	Brutine
P021	Calcum eyeride
P123	Comphene, estechters-
P103	Carteminidectionals and
P022	Carton troutide
PORT.	Carton doubles
P006	i Certainyl chieride

Mezardous wasta No	Substanc +
P033	Chlonne cyenide
P023	Chlorosostaldehyde
P026	p-Chlorosnime 1-(o-Chlorophenyf)thioures
P027	3-Chloropropionitrile
F029	Copper cyandes
P030	Oyerades (soluble oyerade selts), not elec- where specified
P031	Cyanogen
P033	Overagen chloride Dichloropherylerane
P037	Dieldnn
P036	Districtance
P030	O,O-Distryl S-[2-(ethylthic)ethyl] phosphoro- diffusese
P041	Distryl-p-nerophenyl phosphate
P040	O.O-Dietryl O-pyrazmyl phosphorothoose
P043	Discoropyl fluorophospheta Dimethosis
P045	3,3-Dimetryl-1-(metrylthio)-2-butenone. O-
P071	((methylamino)certamy) distrie O,O-Dimethyl O-p-ntrephenyl pheephoro-
	those
P082	Density/introcernine
P046	aiphe, alphe-Dimethylehensetylemine 4,6-Dintro-o-creati and sets
P034	4,6-Dinero-o-cyclohetryfphenol
P044	2.4-Dintrophenol Dinceeb
P086	Diphospheremide, ottomethyl-
P039	Deutoton 2.4-Othioburet
P109	Ottniopyrophosphonic soid, tetraethyl ester
P050	Endocultan
P068	Endothell Endre
P042	Epnephane
P046	Etheremne, 1,1-dimethyl-3-phonyl- Etheremne, N-methyl-N-rutroso-
P101	Ethyl cyanide
P064	Ethyleriane Fernanur
P056	Fuorre
P057	Fluorescetamide Fluorescetta acid, sodium sett
P065	Fulminic scid, mareury(II) self. (FLT)
P050	Heptechtor
P051	1,2,3,4,10,10-Hexastiaro-0,7-apeny- 1,4,4a,5,6,7,8,8a-astatysiro-ando,endo-
	1,4:5,8-dimethanonagnificiens
P037	1,2,3,4,10,10-Hexachtero-0,7-epony- 1,4,4e,5,5,7,8,6e-ectahydro-endo,exo-
	1,4:5,8-demetheronephthesene 1,2,3,4,10,10-Heasthlero-1,4,4e,5,8,8e-
P060	1,2,3,4,10,10-Hemechiero-1,4,4e,5,8,8e- hemerysiro-1,4:5,8-ensis, ensis-dimeth- en-
	cnephthelene
P004	1,2,3,4,10,10-Herechter-1,4,4e,8,8,8-
	heaphydro-1,4:5.5-endo.exio- dmethenenghthelene
P060	dimethenenephthalene Heaschlorphesshydro-eno.eno-
P062	dmetherenephthetene
P116	Housethyl tetrapheephete Hydrapheeshelheephete Hydraphee, methyl-
P060	Hydrogenic acid
P083	. Marchan Cando
P086	Hydragen sheephide leccyanic acid, mothyl ester
P007	.) 3(2H)-leosazzione, 5-(aminomethyl)-
PO01	. Mercury, (acetato-O)phenyl-
P006	Maraury Surrenase (FLT)

Hazardous waste No.	Substance
P112	Methane, letranizo- (R)
P118	Methanethiol, Inchloro-
P060	4,7-Methano-1H-indane, 1,4,5,6,7,8,8-hap-
P086	tachtoro-3e,4,7,7e-tetrahydro- Methomyl
P067	2-Methylazinane
P064	. Methyl hydrazine . Methyl isocyanate
P000	2-Methylactoratile
P071	Methyl perathion
P072	alphe-Naphthylthioures
P073	Nickel cerbonyl Nickel cyensde
P074	Nickel(II) cyanide
P073	Nickel tetracerbonyl
P075	Nicotine and salts
P077	Nitro ande s-Nitrosnitre
P078	Nitrogen dictode
P076	Nitrogen(II) crede
P078	Nitrogen(IV) cade Nitroglycenne (R)
P082	N-Nerosodimethylamine
P084	N-Nitrosometrylvnylemne
P060	5-Nortomene-2,3-dimethenot, 1,4.5.6,7,7-httr- schioro, cyclic suitte
POBS	Octamethylpyrophosphoramide
P087	Osmium celdo
P087	Comum terrorde
P006	7-Outbioyoto (2.2.13haptane-2,3-dicarticaylic acid
P000	Perathion
P034	Phenoi, 2-cycloheryl-4,6-dinitro
P048	Phonoi, 2,4-dintro- Phonoi, 2,4-dintro-6-moltry-
P020	Phanol, 2.4-dintro-8-(1-methyloropyi)-
P006	Phonol, 2,4,6-trintro-, ammonium ant (R)
P094	Phenyl dichlorograme Phenylmerounc accepte
P083	N-PhonyRhoures
P084	Phorate
P006	Phosphie Phosphine
P041	Phosphore and dethi p-nereshani ester
P044	Phosphoro acid, diethyl p-ntrephenyl esser Phosphorodithioic acid, O.O-dimethyl 8-12-
P043	(mothysemino)-2-espethylicetor Phasehorofluenc acid, bm(1-methylethyl)-
F-0-3	atter
P084	Phosphorothicic acid, O.O-disthyl 8-
	(athythap)methyl ceter Phosphorothoca sold, C.O-disthyl O-ip-nitro-
P088	phenyl) color
P040	Prosphorothest sold, O.D-diethyl C- pyreanyl
	Color Colored Colored Colored
P007	Phosphorethes: soid. O.O-dimethyl O-Le-(Id- methylamino)-sulfonyl/phenyl/sester
P110	. Plumpers, weserny.
P000	Potessium syenide Potessium siver cyenide
P070	Propersi. 2-metryl-2-(metrylthio) C-
	((metrytermne)cerbonyl)cethe
P101	Propenentrie
P027	Progenentrie, 3-chloro- Progenentrie, 2-hydrony-2-mathyl-
P061	1,2,3-Propensitiol, trinstrate- (R)
P017	2-Prosenone, 1-bromo-
P102	Propengi alcohol 2-Propensi
P005	2-Propen-1-ol
P067	1.2-Propyterumne

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Hezardous waste No.	Substance
P075	Pyriame, (\$)-3-(1-methyl-2-pyrrolicinyl)-, and sets
P111	Pyrophosphonic acid, termentyl ester
P103	Selencures
P104	Silver cyerate
P105	Sodium azide
P106	Sodum cyanide
P107	Strongum author
P106	Sayonnon-10-one, and salts
P018	Strychnidin-10-one, 2.3-dimethoxy-
P108	Strychrane and salts
P115	Sultuno acid, thelleum(I) salt
P109	Tetraethylosthiopyrophosphale
P110	Tetraethyl lead
P111	Tersettytyrophosphate
P112	Teraneromethene (R)
P062	Terreprosphono acid, hasseshyl ester
P113	Thefic gade
P113	Thelium(III) costs
P114	Thelium(I) selents
P115	Theisum(I) sulface
PO45	Tholenax
P048	Stdeadarana diameta
P014	Thephenol
P116	Thiopinioscopic derive Thiopeniosrbande Thioseniosrbande Thioseniosrbande
P026	Thourse, (2-chiorophenyl)-
P072	Thouse, 1-nechthalenyi-
P083	Thouse, phonyl-
P123	Tomphene
P118	Trichtgromethenethiol
P119	Vanadic acid, emmonium self
P120	Vanegum pentoxide
P120	Vanadium(V) oxide
.001	Wartenn, when present at concentrations
	greater than 0.3%
P121,	
P122	Zinc phosphide (R,T)
P122	Zinc phosphide, when present at concentra-
	tions greater than 10%

(f) The commercial chemical products, manfacturing chemical intermediates, or off-specification commercial chemical products referred to in paragraphs (a) through (d) of this section, are identified as toxic wastes (T), unless otherwise designated and are subject to the small quantity generator exclusion defined in § 261.5 (a) and (g).

[Comment: For the convenience of the regulated community, the primary hazardous properties of these materials have been indicated by the letters T (Toxicity). R (Reactivity). I (Ignitability) and C (Corrosivity). Absence of a letter indicates that the compound is only listed for toxicity.]

These wastes and their corresponding EPA Hazardous Waste Numbers are:

Hazardous Waste No.	Substance
WESTS NO.	
12001	Acetaidefryde (I)
U001 U034	Acetaidehyde, thichioro-
U187	Acetamide, N-(4-ethoxyphenyl)-
U005	Aceterride, N-9H-Ruoren-2-yi-
U112	Acesc acid, ethyl eater (I)
U144	Acetic acid, lead salt
U214	Acetic acid, thefium(i) selt
U002	Acetone (I) Acetonstrie (I,T)
U248	3-(alghe-Acetonylbenzyl)-4-hydroxycoumann
V4	and sets, when present at concentrations
	of 0.3% or less
U004	Acetophenone
U005	2-Acetylemenofluorene
U006	Acetyl chionde (C.Pl.T)
U007	Acrylemide Acrylic acid (1) Acrylonitifie
U009	Acrysic acris (i)
U150	Alenne, 3-(p-bis(2-chlorosthyl)emino)
U 150	phenyl-, L-
U328	2-Americane
U3\$3	4-Ammo i-mathylbensene
U011	America
U012	Anime (I,T)
U014	Auramine
U015	Azzadina
0010	Asimo(2:3:3.4)pymoto(1:2-e)indote-4.7-dione. 6-emino-6-[((aminocarbonyl) snylmethyl]-
	1,1e,2.6,6e,8b-hexalydro-8e-methoxy-5-
	methyl-,
U157	Benz(j)accentrylens, 1,2-dhydro-3-methyl-
U016	Beng(c)acndine
U016	3.4-Serascodine
U017	Bergal chlonde Beng(a)enthracens
U016	1.2.Beconstitutions
U094	1.2-Benzanthracene 1.2-Benzanthracene, 7,12-dimethyl-
U012	Bensenemene (1,T)
U014	Benzenames, 4,4'-aprioremaloy/bis(N,N-d-
	methyl-
U048	Bensenemne, 4-chloro-2-methyl-
U093 U158	Bersenemine, N.N'-dimethyl-4-phenylezo- Bersenemine, 4,4'-methylenetis(2-chloro-
U222	Benzenemene 2-metrot, hydrechlonde
U181	Benzenemene 2-methyl-, hydrachtonde Bengenemene, 2-methyl-5-meto
U019	Benzene (I,T)
U036	Sensoneacosc ecid, 4-chiero-elene-(4-chiero- phenyl)-elene-hydroxy, othyl ester
1	phonyll-alpha-hydraxy, allryl color
U030	Benzens, 1-bromo-4-phenoxy-
U037	Senzene. chloro- 1,2-Benzenedcersoxylic acid anhydride
U190 U026	1,2-Bergenescarbaryic acid. (5002-097)-
	hand) accer
U069	. Concentration of the contract of the contrac
U068	1,2-Benzenedicerboxylic scid, diethyl ester
U102 U107	1,2-Benzenedicarboxylic acid, di-n-octyl deter 1,2-Benzenedicarboxylic acid, di-n-octyl deter
	Senzene, 1,2-dichloro-
U073	Servere 13-defines
U072	Bergene, 1,3-dichloro- Bergene, 1,4-dichloro-
U072	Benzone, (dichloromethyl)-
U223	· Benzone, 1.3-Georgeogenetryl- (R.T)
U239	Benzene, dimethyl-(I,T)
U201	1,3-Bergenediol
U127	Benzene, hexachloro- Benzene, hexachloro- (I)
U056	Benzana, hydroxy-
U220	Bengane, mathyl-
11105	Berrana 1-math4-1-2-4-dintro-
U106	Senzene, 1-metryl-2,6-dratto- Senzene, 1,2-metrylenedoxy-4-ellyl- Senzene, 1,2-metrylenedoxy-4-properyl-
U203	Bensons, 1,2-mothytenedicky-4-ellyl-
U141	J margane, 1,2-methylenedicsy-4-properyl-

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Li Erwa, 1,2-darono	587	Catalanadaryda		á
Character Nation Addition	U174	Company of the Compan		
D-N-propyrations/fire	111	Creases		ğ
ť	0110			
12. Danamendadore		ij		Ş
		Granton		E
2.6-Creatoname	919	Ş		Š
2.4-Orrerotation	200	Chargements matryl other		Ē
Dimetryl eurose	28			S
Directly provides	Ŕ	2-Oriental was offer		
3	C101	1.Charles Samuel Control of the Cont	i	
13-Date of the last		CHORDSHIER		9
	9	Спотвридана	-	8
		Chloridana, technosis		8
3.3 Onesystematics	8	Charambuck		ğ
7.12-Ornery Carata (a) and masons	5	Chloral		ğ
	6	Carsony fluence (R.1)		8
Directorario (1)	5	Carton Managhanda		5
i		Carton Cardinates in 1		3 8
The Contract of the Contract o		3		215
1		Carbanoyi chiones, dimeny-		Ş
1	COM	\$	-	Ę
O.O Cigarys & mustys and appropriate	U087	Carbanida, N-mashy-N-marcao		55
ŀ		Comments Manufacturery, corp. com	İ	
	100			Ş
1.3-Caracroproposition (LT)		Calcum chromate		S
1,3-Ochbropropere	1083	Caccoylic scid		5
1		ž		ğ
2.4-Dichloropheromyscoatc acid, selfs and	38	2-Busine, 1,4-dictions (I,T)		ş
2.8-Dichigrophenoi		2		5
Š	6	2-Buttone partially (R.T.)		
-	1025			
1. Designation				•
	100	Butanosc acid. 4-(Bas/2-chicrosethyl)amino)		8
critoro digitariya d	5080	WAR. N. DURNER		2
bergerade		3 ∴		2
1.5-Dicharo-H-(1,1-dimetryl-2-propyriyi)	U192	4-Bromophenyi phenyi ether		8
5	U075	•		5
ĸ	1074			5
1.3 - Dighteroberodine	C073	-		
-Derivorbergene	L073	ş		
	38	BB(2-CHOROTON)) INSURAN		Ş
\$-(2.3-Canadamy) seast-opperature contract	0012	1		8
Dough provides	5			ã
1.2-Opposito-Generations	LORG	Igneryi).4.4'diamere	-	3
Dibenz(a.i)pyrene	1004	₹.		ğ
1.2:7.8-Oberacpyrene	UOM.	2.2 Bourges (I.T)		
1.2.5.6-Obsergantingcome				5
Diservice of landstream				1
(5.1)	1991	3,4-Benzopyrana		2
	CO12	Benzo(a)pyrene		22
		Benzo(j.k.)fluorene		28
Decacharoccuryoro-1,3,4-matheno-214	U142	1.2-Bergsottspoin-3-ons. 1,1-dioxide		202
	LOS1			D21
000	U080	Bergens, 1.3.5-broke (A.T)		3 6
Dispose	089			N
2 Ad-D same and many	265			8
	0130	Bergeneeufonic acid chloride (C.R)		8
3	U057	e, pentachtoro-nero-	-	185
Cyclonecane (I)	U056	Benzene, peniachoro-		Š
ģ.	U197	Bergera out (17)		
Composin transfer	U246			8
			_	
Substance	Waste No	Substance	Weste No	ž į

25%

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Hazardous Waste No.	Substance	Hazardous Waste No.	Substance
U078	Ethene, 1,1-dichloro-	U140	leobutyl acconci (I,T)
U377	Etnane, 1,2-dichloro-	U141	Isosatrole
J114	1.2-EthanedlyRiscarbemodifhoic acid	U142	Kepons
J131 J024	Ethans, 1,1,1,2,22-hexachioro- Ethans, 1,1'-{methylenebus(cxy)]bus(2-chioro-	U143	Lancarpine Land acetate
,003	Emerantie (I, T)	U145	Leed phosphate
J117	Ethans,1,1'-cayble- (I)	U146	Leed subscripts
J025 J184	Ethens, 1,1'-crybs(2-chloro- Ethens, sentechloro-	U129 U147	Lindene
J208	Ethene, 1,1,1,2-tetrachioro-	U148	Maleic hydrazide
200	Ethene, 1,1,2,2-tetrachioro-	U149	Materiaria
J218	Ethenethicemide	U150	Melphelen
/247	Ethane, 1,1,1,-erchloro-2,2-bis(p-methoxy-	U151	Mercury Methecrylonitrile (I,T)
J227	ghenyi). Estene. 1,1,2-elchioro-	U002	Methanamine, N-methyl- (I)
J043	Ethens, chloro-	U029	Methene, bromo-
J042	Ethens, 2-chlorosthony-	U045	Methens, chloro- (1,T)
J078 J078	Ethens, 1,1-dichloro- Ethens, trans-1,2-dichloro-	U046	Methens, chloromethoxy- Methens, dibromo-
210	Ethens, 1,1,2,2-terachtoro-	U000	Methane, dichloro-
J173	Ethanal, 2,2"-(nitroscrimo)bis-	U075	Methane, dichlorodifluoro-
	Ethenene, 1-phonyl-	U156	Methans, indo-
J006 J366	Ethensyl etterate (C.R.T) 2-Ethenystheriol.	U119 U211	Methanicalloric sold, ethyl either Methanic, tetrachtoro-
U112	Ethyl sesses (f)	U121	Methers, trafsprofugre-
U113	Ethyl assylate (I) Ethyl carbonate (urothen)	U183	Methanethial (1,17)
الحدال	Ethyl carbonate (urethen)	U225	Methene, tribromo-
J036	Ethyl 4.4'-dightersbersslate Ethylene glycol monoethyl ether.	U044 U121	Methane, trichicro- Methane, trichicrofluoro-
U114	Ethylenebuidthucertemic acid)	U123	Methanoic and (C,T)
JO67	Elylene usromide	U036	4,7-Methanoinden, 1,2,4,5,6,7,8,6-eda
J077	Ethylene dichloride		chiere-Se,4,7,7a-tetrelayire-
U116	Ethiene exide (1,T) Ethylene tricures	U154	Methenoi (1) Methepyrtiene
U117	Ethyl other (I)	U247	Methanychior.
U078	Ethylidene dehlande	U154	Mathyl aleshol (I)
U118	Ethylmethecrylete	UC29	Mothyl bronucie
U118 U1 36	Etryl metheries.ifonete Ferrie destran	U186	1-Methythutediene (f) Methyt ettande (f.T)
U120	Fluorenthene	U156	Methyl chlorocerbonete (I,T)
U122	Fermaldehyde	U226	Methylchlorolom
U123	Forms and (C.T)	U187	3-Methylcholenthrene 4.4'-Methylenebus(2-shipmentline)
U124	Furan (i) 2-Furancerisconsiderlysis (i)	U192	2.2 Medical annual A. Salahian annual
U147	2.5-Furundone	U086	2,2"-Motivionable(3,4,6-bishisrephanel) Methylane brainide
U213	Furnit telephone (I)	U090	Mathylana eristida
U125 U124	Furtural (f)	U122	Matrijtana estala Matrijtana estala katana (LT)
U206	Furturen (f) D-Ghusspyranses, 2-desmy-2(3-methyl-3-retro-	U160	Mothyl stryl hatens parends (FLT)
	decreedo)-	U136	Marrie Indida
U126	Glyandytanantyda	U161	Mothyl testsulyf beterno (I) Mothyl methenylate (I,T)
U1 63	Guerndine, N-nitrese N-methyl-N'nitro- Heisenterseene	U163	N-Methyl methetrytete (I,T) N-Methyl-M'-retro-M-retroseguerádno
U126	Henchierstutedene	U161	4-Methyl-2-pentanens (I)
J129	Henselterseyclehaugne (gartette isomer)	U164	Methythiouract
U130	Hamahiaraayalapartadlaha	U010	Milenyon C
U131 U132	Homestorestiere Homestorestiere	U069	5,12-Neghthasenedene, (86-ds) 6-asstyl-10 (3-amino-2,3,6-trideoxy-alpha-L-hmo-
U243	Hemathersersens		hemopyranosyrjenyl3-7,8.8,10-tetrahydro-
U133	Hydramne (FLT)		6,8,11-trilydramy-1-matheny-
<u> </u>	Hydrame, 1,8-detry-	U105	Naphthalana Naphthalana, 2-ahlara-
J000	Hydrama, 1,1-dinatry- Hydrama, 1,2-dinatry-	U047	1,4-haptitusenedene
U100	Historia, 1,3-denoni-	U236	2,7-Neghtheranecoullaric said. 3.5'-(3.5'-4
J134	Hydrofluoric and (C.T)		mathyl-(1,1'-baptarryl)-4,4'dyl)}-bas
U134	Hydregen Suchée (C,T)		(aso)sis(E-emm-4-hydrany)-Johnsodum
U1 36	Hydragen authos Hydragerause, 1-methyl-1-phenylethyl-(R)	U106	1,4,Neshtheeuinone
J596 J196	Hydranydmethylarane exce	U167	1-Neutrino
J116	2-bridgestidnethone	U166	2-Neghthylemma
J137	indene(1.2.3-ed)pyrane	U167	estro-Negitalnylemine Inde-Negitalnylemine
U139	J train dispiran	U166	Late-Healthylamure

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Hezardous Waste No.	Substance	Hazardous Waste No.	Substance
JC26	2-Nachthylamine, N,N'-bis(2-chioromethyl)-	U191	Pyndine, 2-methyl-
J169	Nerobenzene (I,T)	U164	4(1H)-Pynmonone. 2,3-dihydro-6-methyl-2-
1170	p-N/trophenol		themo-
171	2-Noropropene (I,T)	U180 U200	Pyrrole, tetratrydro-N-retroso- Reserpine
172 173	N-Nérosco-n-butylemne N-Nérosco-therolemne	U201	Resorance
74	N-Nitrosociethylamine	U202	Seothern and selts
11	N-Netroso-N-propytemine	U203	Setrole
6	N-Nitroso-N-ethylures	U204	Selenious acid
77	N-Nitroso-N-methytures	U204	Selentum dioxide
78	N-Nation-N-methylurethane	U205 U015	Selentum dieutfide (R,T)
79	N-Nitrosopipendine	See F027	L-Senne, diszoscetate (ester)
80 81	N-Nitrosopymolidine 5-Nitro-o-lokadine	U009	4,4'-Silbenedici, alphe alphe'-diethyl-
93	1,2-Oxatholene, 2,2-dioxede	U206	Streptozolocin
58	2H-1,3,2-Oxezephosphonne, 2-(bis(2-chioro-	U135	Sulfur hydride
	ethyllemino litetratydro-, carde 2-	U103	Sulfuric ecid, dimethyl easer
15	Ourane (I,T)	U189	Suffer phosphide (R)
41	Ourses, 2-(chloromethyl)-	See F027	Sulfur selerade (R,T) 2.4.5-T
82 83	Persidenyde Persidenyde	U207	1.2.4.5-Tetrachierobergene
M	Pertachiaronhane	U206	1,1,1,2-Tetrachiorosthene
85	Pentachiprontrobertsene	U200	1,1,2,2-Tetrachiorosthane
● F027	Pentechtorophenol	U210	Tetrachioroethylene
4	1,3-Pentachane (I)	See F027 U213	2,3,4,6-Tetrachiorophenol
87	Phonecoun	U214	Tetrahydroturan (f) Thelium(i) ecosse
₩ ₩	Phenol Phenol, 2-chioro-	U215	Theilum(i) carbonate
30	Phenol, 4-chloro-3-methyl-	U218	Theillum(I) chlonde
6 1	Phenol, 2.4-dichloro-	U217	Theffum(I) nersee
62	Phenol, 2.6-dichloro-	U218	Thiosostemide
01	Phenoi, 2,4-dimetryl-	U1 53	Thomstend (1,T)
70	Phenol, 4-nitro-	U219	Thouses Three
• F027 	Phenol, pentachioro- Phenol, 2,3,4,6-tetrachioro-	U220	Tourne
Do	Phenoi, 2,4,5-inchioro-	U221	Toluenedemine
Do	Phenol, 2.4.5-Inchloro-	U223	Toluene discoplanate (R.T)
7	1,10-(1,2-phenylene)pyrene	U329	a-Toludine
<u> </u>	Phosphonic scid. Lead selt	U222	O-Tolusine hydrachlande o-Tolusine
87	Phosphorodithioic acid, 0,0-diethyl-, 8-methy-	U011	1H-1.2.4-Trigger-3-emine
	Phosphorous sulfide (R)	U226	1,1,1-Trightoroethane
·····	Phthalic anniverse	U227	1,1,2-Trientorosthene
D1	2-Propine	U226	Trichloreethene
92	Pronemide	U220	Triphiarosthylene
<u> </u>	1-Proponerune (I,T)	U121	Trionisremenofusremethene 2.4.5-Trionisrophenol
10 8 6	1-Propenemne, N-propyl- (I) Propene, 1,2-dibramo-3-chtoro-	Do	2.4.6-Trichlorophenol
49	Propensioninis	Do	2.4,5-Trightorophenoxyeostic acid
71	Propens, 2-nero- (1,T)	U234	sym-Trimestersone (R,T)
27	Propens, 2,2'onybis[2-chloro-	U182	1,3,5-Triemane, 2,4,5-threetyl-
9 3	1,3-Propone autone	U236	Tre(2,3-ditremepropyl) phosphele Trypen blue
35 26	1-Properol, 2.3-dibrono-, pheephese (3:1)	U237	Uraci, Sibio(2-chierumethytemmo)-
40	1-Propend, 2.3-econy- 1-Propend, 2-methyl- (I,T)	U237	Uraci mustard
02	2-Progenone (I)	U043	Vinyl chlends
207	2-Propenantide	U248	Wartern, when present at consumptions of
84	Propens, 1,3-dichtoro-	11000	0.3% or less
43	1-Propene, 1,1,2,3,3,3-hexachioro-	U239	Xylens (f) Yohmber-16-cersonylic edid, 11,17-dimeth
S2	2-Propenentine	V200	cay-18-((3.4.5-timetany-barasy))any)
06	2-Proponentitie, 2-methyl- (1,17) 2-Proponenc east (f)		mothyl cater
12	2-Propencia and, othyl ester (f)	U248	Zinc phosphide, when present at esnounts
18	2-Propencic acid, 2-methyl-, othyl aster		sons of 10% or less.
L2	2-Propensic acid, 2-methyl-, methyl ester (I,T)		<u> </u>
P027	Propients acid, 2-(2,4,5-trichloropnenczy)-		
<u> </u>	n-Propytemene (LT)		by the Office of Management
ğ	Propyrane dichionile Pyridine		set under control number 2050-
<u> </u>	Pyridine, 8-[(2-(dimethytemeno)-2-thenyte-	0047)	
	minol-	[45 FR '	78529, 78541, Nov. 25, 1980, a

Appendix II

Subpart C-Characteristics of Hezerdous Waste

\$ 261.20 General.

(a) A solid waste, as defined in § 261.2, which is not excluded from regulation as a hazardous waste under § 261.4(b), is a hazardous waste if it exhibits any of the characteristics identified in this subpart.

(Comment: § 262.11 of this chapter sets forth the generator's responsibility to determine whether his waste exhibits one or more of the characteristics identified in this subpart1

(b) A hazardous waste which is identified by a characteristic in this sub-part, but is not listed as a hazardous waste in Subpart D. is assigned the EPA Hazardous Waste Number set forth in the respective characteristic in this subpart. This number must be used in complying with the notification requirements of section 3010 of the Act and certain recordkeeping and reporting requirements under Parts 262 through 265 and Part 270 of this chapter.

(c) For purposes of this subpart, the Administrator will consider a sample obtained using any of the applicable sampling methods specified in Appendix I to be a representative sample within the meaning of Part 260 of this

chapter.

[Comment: Since the Appendix I sampling methods are not being formally adopted by the Administrator, a person who desires to employ an alternative sampling method is not required to demonstrate the equivalency of his method under the procedures set forth in \$5,250,20 and 250,21.1

[45 FR 33119, May 19, 1980, as amended at 48 FR 14294, Apr. 1, 1983)

261.21 Characteristic of ignitability.

(a) A solid waste exhibits the characteristic of ignitability if a representative sample of the waste has any of

the following properties:

(1) It is a liquid, other than an aqueous solution containing less than 24 percent alcohol by volume and has flash point less than 60°C (140°F), as determined by a Pensky-Martens Closed Cup Tester, using the test method specified in ASTM Standard D-93-79 or D-93-80 (incorporated by reference, see § 260.11), or a Setaflash Closed Cup Tester, using the test method specified in ASTM Standard D-3278-78 (incorporated by reference, see § 260.11), or as determined by an equivalent test method approved by the Administrator under procedures set forth in §§ 260.20 and 260.21.

(2) It is not a liquid and is capable, under standard temperature and pressure, of causing fire through friction, absorption of moisture or spontaneous chemical changes and, when ignited, burns so vigorously and persistently that it creates a hazard.

(3) It is an ignitable compressed gas as defined in 49 CFR 173.300 and as determined by the test methods described in that regulation or equivalent test methods approved by the Administrator under \$\$ 260.20 and 260.21.

(4) It is an oxidizer as defined in 49

CFR 173.151.

(b) A solid waste that exhibits the characteristic of ignitability, but is not listed as a hazardous waste in Subpart D, has the EPA Hazardous Waste Number of D001.

[45 FR 33119, May 19, 1980, as amended at 46 FR 35247, July 7, 1981]

\$ 261.22 Characteristic of corrosivity.

(a) A solid waste exhibits the characteristic of corrosivity if a representative sample of the waste has either of the following properties:

- (1) It is aqueous and has a pH less than or equal to 2 or greater than or equal to 12.5, as determined by a pH meter using either an EPA test method or an equivalent test method approved by the Administrator under the procedures set forth in \$\$ 260.20 and 260.21. The EPA test method for pH is specified as Method 5.2 in "Test Methods for the Evaluation of Solid Waste, Physical/Chemical Methods" (incorporated by reference. 4 260.11).
- (2) It is a liquid and corrodes steel (SAE 1020) at a rate greater than 6.35 mm (0.250 inch) per year at a test temperature of 55°C (130°F) as determined by the test method specified in NACE (National Association of Corrosion Engineers) Standard TM-01-69 as standardized in "Test Methods for the Evaluation of Solid Waste, Physical/ Chemical Methods" (incorporated by reference, see § 260.11) or an equivalent test method approved by the Administrator under the procedures set forth in \$\$ 260.20 and 260.21.

(b) A solid waste that exhibits the characteristic of corrosivity, but is not listed as a hazardous waste in Subpart D, has the EPA Hazardous Waste Number of D002.

[45 FR 33119, May 19, 1960, as amended at 46 FR 35247, July 7, 1981]

\$ 261.23 Characteristic of reactivity.

(a) A solid waste exhibits the characteristic of reactivity if a representative sample of the waste has any of the following properties:

(1) It is normally unstable and readily undergoes violent change without detonating.

(2) It reacts violently with water.

(3) It forms potentially explosive mixtures with water.

(4) When mixed with water, it generates toxic gases, vapors or fumes in a quantity sufficient to present a danger to human health or the environment.

(5) It is a cyanide or sulfide bearing waste which, when exposed to pH conditions between 2 and 12.5, can generate toxic gases, vapors or fumes in a quantity sufficient to present a danger to human health or the environment.

(6) It is capable of detonation or explosive reaction if it is subjected to a strong initiating source or if heated

under confinement.

(7) It is readily capable of detonation or explosive decomposition or reaction at standard temperature and pressure.

(8) It is a forbidden explosive as defined in 49 CFR 173.51, or a Class A explosive as defined in 49 CFR 173.53 or a Class B explosive as defined in 49 CFR 173.88.

(b) A solid waste that exhibits the characteristic of reactivity, but is not listed as a hazardous waste in Subpart D, has the EPA Hazardous Waste Number of D003.

\$ 261.24 Characteristic of EP toxicity.

(a) A solid waste exhibits the characteristic of EP toxicity if, using the test methods described in Appendix II or equivalent methods approved by the Administrator under the procedures set forth in \$\$ 260.20 and 260.21, the extract from a representative sample of the waste contains any of the contaminants listed in Table I at a concentration equal to or greater than the respective value given in that Table. Where the waste contains less than 0.5 percent filterable solids, the waste itself, after filtering, is considered to be the extract for the purposes of this section.

(b) A solid waste that exhibits the characteristic of EP toxicity, but is not listed as a hazardous waste in Subpart D. has the EPA Hazardous Waste Number specified in Table I which corresponds to the toxic contaminant causing it to be hazardous.

TABLE !- MAXIMUM CONCENTRATION OF CON-TAMINANTS FOR CHARACTERISTIC OF EP Toxicity

EPA hazerdous waste number	Contaminent	Maximum concentra- tion (miligrams per ster)
D004	Arsenic	5.0
D005	Barum	100.0
	Cadmum	1.0
	Chromum	5.0
	Lend	5.0
D009		0.2
D010		1.0
D011	Siver	5.0

APPENDIX I—REPRESENTATIVE SAMPLING METRODS

The methods and equipment used for sampling waste materials will vary with the form and consistency of the waste materials to be sampled. Samples collected using the sampling protocols listed below, for sam-pling waste with properties similar to the indicated materials, will be considered by the Agency to be representative of the waste.

Extremely viscous liquid—ASTM Standard D140-70 Crushed or powdered material— ASTM Standard D346-75 Soil or rock-like material—ASTM Standard D420-69 Soillike material—ASTM Standard D1452-65

Ply Ash-like material—ASTM Standard D2234-76 [ASTM Standards are available from ASTM, 1916 Race St., Philadelphia. PA 191033

PA 19103)
Containerised liquid wastes—"COLIWASA"
described in "Test Methods for the Evaluation of Solid Waste, Physical/Chemical
Methods," " U.S. Environmental Protection Agency, Office of Solid Waste, Washington, D.C. 20460. [Copies may be obtained from Solid Waste Information, U.S. Environmental Protection Agency, 28 W.

Environmental Protection Agency, as w. St. Clair St., Cincinnati, Ohio 452681
Liquid waste in pits, ponds, lagoous, and similar reservoirs.—"Pond Sampler" described in "Tast Methods for the Evaluation of Solid Waste, Physical/Chemical

This manual also contains additional information on application of these protocols.

AFFENDIX II-EP TOXICITY TEST PROCEDURES

A. Extraction Procedure (EP)

1. A representative sample of the waste to be tested (minimum size 100 grams) shall be obtained using the methods specified in Appendix I or any other method capable of yielding a representative sample within the meaning of Part 260. (For detailed guidance on conducting the various aspects of the EP see "Test Methods for the Evaluation of Solid Waste, Physical/Chemical Methods" (incorporated by reference, see § 260.11).]

2. The sample shall be separated into its component liquid and solid phases using the method described in "Separation Proce-dure" below. If the solid residue a obtained using this method totals less than 0.5% of the original weight of the waste, the residue can be discarded and the operator shall treat the liquid phase as the extract and proceed immediately to Step 8.

3. The solid material obtained from the Separation Procedure shall be evaluated for its particle size. If the solid material has a surface area per gram of material equal to, or greater than, 3.1 cm² or passes through a 9.5 mm (0.375 inch) standard sieve, the operator shall proceed to Step 4. If the surface area is smaller or the particle size larger than specified above, the solid material shall be prepared for extraction by crushing, cutting or grinding the material so that it passes through a 9.5 mm (0.375 inch) sieve or, if the material is in a single piece, by subjecting the material to the "Structural Integrity Procedure" described below.
4. The solid material obtained in Step 3

shall be weighed and placed in an extra with 16 times its weight of deionised water.

Do not allow the material to dry prior to weighing. For purposes of this test . An ac ceptable extractor is one which will impart sufficient agitation to the mixture to not only prevent stratification of the sample and extraction fluid but also insure that all sample surfaces are continuously brought into contact with well mixed extraction

5. After the solid material and deionized water are placed in the extractor, the operator shall begin agitation and measure the pH of the solution in the extractor. If the pH is greater than \$.0, the pH of the solution shall be decreased to 5.0 ± 0.2 by adding 0.5 N scetic acid. If the pH is equal to or less than 5.0, no acetic acid should be added. The pH of the solution shall be monitered, as described below, during the course of the extraction and if the pH rises above 5.2, 0.5N acetic acid shall be added to bring the pH down to 5.0 \pm 0.2. However, in no event shall the aggregrate amount of acid added to the solution exceed 4 ml of acid per gram of solid. The mixture shall be agitated for 24 hours and maintained at 20 40°C (68'-104'F) during this time. It is recommended that the operator monitor and adjust the pH during the course of the extraction with a device such as the Type 45-A pH Controller manufactured by Chemtrix. Inc., Hillsboro, Oregon 97123 or its equivalent, in conjunction with a metering pump and reservoir of 0.5N scetic acid. If such a system is not available, the following manual procedure shall be employed:

(a) A pH meter shall be calibrated in ac-cordance with the manufacturer's specifica-

(b) The pH of the solution shall be checked and if necessary, 0.5N acetic acid shall be manually added to the extractor until the pH reaches 5.0 ± 0.2 . The pH of the solution shall be adjusted at 15, 20 and 60 minute intervals, moving to the next longer interval if the pH does not have to be adjusted more than 0.5N pH units.

(c) The adjustment procedure shall be

continued for at least 6 hours.

(d) If at the end of the 24-hour extraction period, the pH of the solution is not below 5.2 and the maximum amount of scid (4 ml per gram of solids) has not been added, the pH shall be adjusted to 5.0 ± 0.2 and the extraction continued for an additional four hours, during which the pH shall be adjusted at one hour intervals.

 At the end of the 24 hour extraction period, deionized water shall be added to the extractor in an amount determined by the following equation: V=(20)(W)-16(W)-A

V=ml deionised water to be added

W-weight in grams of solid charged to ex-

A=ml of 0.5N acetic acid added during extraction

7. The material in the extractor shall be separated into its component liquid and solid phases as described under "Separation Procedure."

8. The liquids resulting from Steps 2 and 7 shall be combined. This combined liquid (or the waste itself if it has less than % percent solids, as noted in Step 2) is the extra shall be analyzed for the presence of any of the contaminants specified in Table I of § 261.24 using the Analytical Procedures designated below.

Separation Procedure

Equipment: A filter holder, designed for filtration media having a nominal pore size of 0.45 micrometers and capable of applying a 5.3 kg/cm² (75 psi) hydrostatic pre seure to the solution being filtered, shall be used. For mixtures containing nonabsorptive solids, where separation can be effected without imposing a 5.3 kg/cm³ pressure dif-ferential, vacuum filters employing a 0.45 micrometers filter media can be used. (For

Manardous Waste Streams," EPA 600/2-80-018, January 1960.

The percent solids is determined by drying the filter pad at 80°C until it reaches constant weight and then calculating the percent solids using the following equation:
Percent solids =

(weight of pad + sold) - (tere weight of pad)

- ×100 inited would at named

further guidance on filtration equipment or procedures see "Test Methods for Evaluating Solid Waste, Physical/Chemical Methincorporated by reference, [250.11). Proceedure:

(i) Following manufacturer's directions, the filter unit shall be assembled with a filter bed consisting of a 0.45 micrometer filter membrane. For difficult or slow to filter mixtures a prefilter bed consisting of the following prefilters in increasing pore size (0.65 micrometer membrane, fine glass fiber prefilter, and coarse glass fiber prefilter) can be used.

(ii) The waste shall be poured into the fil-

tration unit.
(iii) The reservoir shall be slowly pressurized until liquid begins to flow from the fil-trate outlet at which point the pressure in the filter shall be immediately lowered to 10-15 psig. Filtration shall be continued until liquid flow ceases.

(iv) The pressure snall be increased step-wise in 10 psi increments to 75 psis and fil-tration continued until flow ceases or the pressurizing gas begins to exit from the fil-

trate outlet.

(v) The filter unit shall be depressurized. the solid material removed and weighed and then transferred to the extraction appara-tus, or, in the case of final filtration prior to analysis, discarded. Do not allow the materi-

This procedure is intended to result in separation of the "free" liquid portion of the waste from any solid matter having a particle size >0.45 μm. If the sample will not filter, various other separation techniques can be used to sid in the filtration. As described above, pressure filtration is employed to speed up the filtration process. This does not alter the nature of the separation. If liquid does not separate during filtration, the waste can be omtrifuged. If separation occurs during centrifugation, the liquid portion (centrifugate) is filtered through the 0.45 µm filter prior to becoming mixed with the liquid portion of the waste obtained from the initial filtration. Any ma-terial that will not pass through the filter after centrifugation is considered a solid and is extracted.

al retained on the filter pad to dry prior to weighing.

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(vi) The liquid phase shall be stored at 4°C for subsequent use in Step 8.

B. Structural Integrity Procedure

Equipment: A Structural Integrity Tester having a 3.18 cm (1.25 in.) diameter hammer weighing 0.33 kg (0.73 lbs.) and having a free fall of 15.24 cm (6 in.) shall be used. This device is available from Associated Design and Manufacturing Company, Alexandria, VA 22314, as Part No. 125, or it may be fabricated to meet the specifications shown in Figure 1.

Procedure

1. The sample holder shall be filled with the material to be tested. If the sample of waste is a large monolithic block, a portion shall be cut from the block having the dimensions of a 3.3 cm (1.3 in.) diameter x 7.1 cm (2.8 in.) cylinder. For a fixated waste, samples may be cast in the form of a 3.3 cm (1.3 in.) diameter x 7.1 cm (2.8 in.) cylinder for purposes of conducting this test. In such cases, the waste may be allowed to cure for 30 days prior to further testing.

2. The sample holder shall be placed into the Structural Integrity Tester, then the hammer shall be raised to its maximum height and dropped. This shall be repeated

fifteen times.

3. The material shall be removed from the sample holder, weighed, and transferred to the extraction apparatus for extraction.

Analytical Procedures for Analyzing Extract Contominants

The test methods for analyzing the ex-

tract are as follows:

1. For amenic, barium, cadmium, chromium, lead, mercury, selenium, silver, endrin. lindane, methoxychior, toxaphena, 2.4. D[2,4-dichlorophenoxyscetic acid] or 2,4,5-TP [2,4,5-trichlorophenoxypropionic acid]:
"Test Methods for the Evaluation of Solid Waste, Physical/Chemical Methods" (incorporated by reference, see § 280.11).

2. (Reserved)
For all analyses, the methods of standard addition shall be used for quantification of species concentration.